

Specific recognition of two MAX effectors by integrated HMA domains in plant immune receptors involves distinct binding surfaces

Liwei Guo^{a,b,1}, Stella Cesari^{c,1}, Karine de Guillen^{d,1}, Véronique Chalvon^c, Léa Mammri^d, Mengqi Ma^{a,b}, Isabelle Meusnier^c, François Bonnot^{c,e}, André Padilla^d, You-Liang Peng^{a,b}, Junfeng Liu^{a,b,2}, and Thomas Kroj^{c,2}

^aState Key Laboratory of Agrobiotechnology, China Agricultural University, 100083 Beijing, People's Republic of China; ^bMinistry of Agriculture and Rural Affairs, Key Laboratory of Pest Monitoring and Green Management, China Agricultural University, 100083 Beijing, People's Republic of China; ^cBGPI, Université de Montpellier, CIRAD, INRA, Montpellier SupAgro, F-34398 Montpellier, France; ^dCBS, Université de Montpellier, CNRS, INSERM, 34090 Montpellier, France; and ^eCIRAD, UMR BGPI, F-34398 Montpellier, France

Edited by Jeffery L. Dangl, University of North Carolina, Chapel Hill, NC, and approved September 26, 2018 (received for review June 21, 2018)

The structurally conserved but sequence-unrelated MAX (*Magnaporthe oryzae* avirulence and ToxB-like) effectors AVR1-CO39 and AVR-PikD from the blast fungus *M. oryzae* are recognized by the rice nucleotide-binding domain and leucine-rich repeat proteins (NLRs) RGA5 and Pikp-1, respectively. This involves, in both cases, direct interaction of the effector with a heavy metal-associated (HMA) integrated domain (ID) in the NLR. Here, we solved the crystal structures of a C-terminal fragment of RGA5 carrying the HMA ID (RGA5_S), alone, and in complex with AVR1-CO39 and compared it to the structure of the Pikp1_{HMA}/AVR-PikD complex. In both complexes, HMA ID/MAX effector interactions involve antiparallel alignment of β -sheets from each partner. However, effector-binding occurs at different surfaces in Pikp1_{HMA} and RGA5_{HMA}, indicating that these interactions evolved independently by convergence of these two MAX effectors to the same type of plant target proteins. Interestingly, the effector-binding surface in RGA5_{HMA} overlaps with the surface that mediates RGA5_{HMA} self-interaction. Mutations in the HMA-binding interface of AVR1-CO39 perturb RGA5_{HMA}-binding, in vitro and in vivo, and affect the recognition of *M. oryzae* in a rice cultivar containing *Pi-CO39*. Our study provides detailed insight into the mechanisms of effector recognition by NLRs, which has substantial implications for future engineering of NLRs to expand their recognition specificities. In addition, we propose, as a hypothesis for the understanding of effector diversity, that in the structurally conserved MAX effectors the molecular mechanism of host target protein-binding is conserved rather than the host target proteins themselves.

plant immunity | immune receptor | rice | *Magnaporthe oryzae* | effector

Intracellular nucleotide-binding domain and leucine-rich repeat protein (NLR) receptors are key elements of plant immunity. They are characterized by a central nucleotide-binding (NB-ARC) and a C-terminal leucine-rich repeat (LRR) domain (1–3). Plant NLRs detect pathogens by recognizing virulence effectors inside host cells and activate defense responses and immunity. Since they confer immunity against many crop diseases, which represent a major problem in agriculture, NLR-coding genes are widely used in crop resistance breeding.

The specific recognition of pathogen effectors by NLRs relies on several mechanisms (4). Some NLRs directly interact with effectors, while others recognize the effector-mediated modification of a host protein that is either a virulence target of the effector (guardee) or a mimic of such a target (decoy) (5, 6). Decoys are thought to stem from the duplication of virulence target genes and to have lost their original cellular function to only serve as effector traps. Recently, an integrated decoy model for effector recognition has been proposed for NLRs carrying noncanonical integrated domains (IDs) and relying on a second, genetically clustered NLR (7). This mechanism was confirmed for *Arabidopsis thaliana* RRS1 that recognizes, through its WRKY

ID, the bacterial effector PopP2, which targets plant WRKY transcription factors (8, 9).

In rice, the paired NLR genes *RGA4* and *RGA5* confer resistance to *Magnaporthe oryzae* isolates carrying the effector genes *AVR1-CO39* or *AVR-Pia*, while *Pikp-1* and *Pikp-2* recognize isolates expressing *AVR-PikD* (10, 11). AVR1-CO39, AVR-Pia, and AVR-PikD are sequence-unrelated, but possess highly similar β -sandwich structures characteristic of the *M. oryzae* AVR and ToxB-like (MAX) effector family in plant pathogenic Ascomycete fungi that has specifically expanded in *M. oryzae* (12, 13). Both RGA5 and Pik-1 contain a heavy metal-associated ID (HMA, also called the Related to yeast ATX1 or RATX1 domain) that is crucial for specific effector recognition through direct binding (10, 13). Structure–function analyses provided detailed insight into Pikp1_{HMA}/AVR-PikD and RGA5_{HMA}/AVR-Pia-binding and established a causal link between these interactions and recognition specificities (13–15).

To further investigate the recognition of MAX effectors by HMA IDs in rice NLRs, we have resolved and functionally validated the crystal structures of RGA5_S (RGA5_{982–1116}), a

Significance

In this study, we provide insight into the mechanism of effector recognition by plant nucleotide-binding domain and leucine-rich repeat proteins (NLRs), a ubiquitous class of immune receptors that plays a central role in crop protection. By structural and functional analysis of a complex between a fungal effector and an integrated decoy domain (ID) from a rice NLR, we demonstrate the importance of IDs in effector recognition and generate crucial knowledge for future engineering of NLRs to expand their recognition specificities. In addition, we propose, as a hypothesis regarding the diversity of fungal effectors, that in structurally conserved effector families, the molecular mechanisms of host target protein-binding are conserved but not the host target proteins themselves.

Author contributions: S.C., A.P., Y.-L.P., J.L., and T.K. designed research; L.G., S.C., K.d.G., V.C., L.M., M.M., and I.M. performed research; L.G., S.C., K.d.G., M.M., I.M., F.B., A.P., J.L., and T.K. analyzed data; and L.G., S.C., A.P., Y.-L.P., J.L., and T.K. wrote the paper.

The authors declare no conflict of interest.

This article is a PNAS Direct Submission.

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Data deposition: The atomic coordinates and structure factors have been deposited in the Protein Data Bank, www.rcsb.org (PDB ID codes 5ZNG and 5ZNE).

¹L.G., S.C., and K.d.G. contributed equally to this work.

²To whom correspondence may be addressed. Email: jliu@cau.edu.cn or thomas.kroj@inra.fr.

This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10.1073/pnas.1810705115/-DCSupplemental.

Published online October 24, 2018.

small C-terminal fragment of RGA5 carrying the HMA domain (*SI Appendix, Fig. S1*), alone and in complex with AVR1-CO39. This shows that specific recognition of the two MAX effectors, AVR1-CO39 and AVR-PikD, by the HMA-IDs of two different rice NLRs, RGA5 and Pikp-1, involves distinct binding surfaces in both partners of the complex. These findings have substantial implications for future engineering of immune receptors carrying HMA IDs to expand their effector recognition specificities.

Results

Previously, C-terminal fragments of RGA5 carrying the HMA ID have been shown to self-interact and to interact with AVR1-CO39 in yeast (10, 16). ITC (isothermal titration calorimetry) measurements with recombinant RGA5_{982–1116} (RGA5_S) and AVR1-CO39 with its signal peptide deleted (dSP-AVR1-CO39) (*SI Appendix, Fig. S1*) showed that their interaction relies on direct binding with an equilibrium dissociation constant (K_d) of 5.4 μ M (17). This K_d is on the same order of magnitude as the one measured for the RGA5_S/dSP-AVR-Pia complex (K_d = 1.8 μ M) (*SI Appendix, Fig. S24*). To reveal the molecular and structural details of AVR1-CO39-binding to the HMA ID of RGA5, we generated crystals for RGA5_S and RGA5_S/AVR1-CO39 diffracting at 1.78 and 2.19 Å resolutions, respectively (17, 18).

RGA5_{HMA} Domain Has a Canonical HMA Fold and Forms a Dimer. The RGA5_S structure was determined by molecular replacement using Phaser (19) with the homologous structure of the copper-binding domain of HMA7 (3DXS) as a search model. This provided a model (PDB accession 5ZNE) containing 73 residues (997–1069). They adopt, as expected, an HMA domain fold characterized by two α -helices and a four-stranded antiparallel β -sheet (topology β_1 - α A- β_2 - β_3 - α B- β_4) (Fig. 1*A* and *SI Appendix, Table S1*). This structure is highly similar to that of the Pikp-1 HMA domain [PDB: 5A6P; sequence identity: 55.1%, rmsd 0.90 Å for 66 (997–1007 and 1010–1064) of 73 residues (997–1069) aligned by Coot (20)] and other HMAs from various organisms (*SI Appendix, Fig. S3 A and B*). The metal-binding motif, characteristic of HMAs, is degenerated in RGA5_{HMA}, as only the first Cys is conserved, while the second is replaced by a Ser.

RGA5_S monomers dimerize through an intimate binding interface that buries 11% of the exposed surface area of RGA5_{HMA} (492 of 4,600 Å²) and that is formed by the secondary structure elements α A, β_2 , and the loop linking them. Binding occurs through hydrophobic interactions and 10 hydrogen bonds distributed on three interaction sites (Fig. 1*A* and *SI Appendix, Fig. S44*). The first site is characterized by the antiparallel alignment of the β_2 s of the two RGA5_{HMA} molecules and leads

to the formation of one continuous β -sheet comprising the two four-stranded β -sheets of the monomers (Fig. 1*A* and *SI Appendix, Fig. S44*). The other two sites are formed by the antiparallel alignment of the α As and by the α A/ β_2 turns of the two RGA5_{HMA} molecules.

A Single Interface Within RGA5_{HMA} Mediates AVR1-CO39-Binding and Self-Interaction. Diffraction data obtained with RGA5_S/dSP-AVR1-CO39 crystals (17) were used to solve the structure of the RGA5_S/dSP-AVR1-CO39 complex using molecular replacement of the RGA5_S structure and the NMR structure of dSP-AVR1-CO39 (PDB: 2MYV). The final refined model contained residues 993–1069 of RGA5 and residues 23–81 of AVR1-CO39 (PDB accession 5ZNG, Fig. 1*B*). The protein structures in the complex are highly similar to the structures of the isolated proteins (RMSD for backbone atoms 0.62 and 1.2 Å, residue range 997–1069 and 23–81, for the RGA5_S and AVR1-CO39, respectively). RGA5_{HMA} and dSP-AVR1-CO39 form a complex with a 1:1 stoichiometry where β_2 of AVR1-CO39 is aligned in an antiparallel manner to β_2 of RGA5_{HMA}, resulting in a continuous antiparallel β -sheet comprising the four-stranded β -sheet of RGA5_{HMA} and β_1 , 2, and 6 of AVR1-CO39 (Fig. 1*B*). Complex stability is mediated by an intimate binding interface that buries 12% and 10%, respectively, of the solvent-exposed surface area of the effector (495 Å²) and RGA5_{HMA} (455 Å²) and that is dominated by hydrophobic interactions. Hydrogen bonds, involving residues of AVR1-CO39, mainly from β_2 and the N terminus, and residues in RGA5_{HMA}, mainly from α A and β_2 , are distributed on three binding areas. (Fig. 1*B* and *SI Appendix, Fig. S4B*).

The first important binding area is formed by W23, K24, and T41 of AVR1-CO39 and V1025 and D1026 of RGA5 and dominated by main- and side-chain hydrogen-bonding between AVR1-CO39^{T41} and RGA5^{D1026} (*SI Appendix, Figs. S4B and S5A*). The second area involves N38 and I39 of AVR1-CO39 and V1028 and E1029 of RGA5 (*SI Appendix, Fig. S5B*). The third interaction site is centered on R1012 and I1030 of RGA5 and D35, V36, N37 of AVR1-CO39 (*SI Appendix, Fig. S5C*).

The binding of AVR1-CO39 to RGA5_{HMA} involves the same interface and many of the same residues as the RGA5_{HMA} homodimerization, suggesting that binding of AVR1-CO39 to RGA5_S competes with RGA5_{HMA} self-interaction and explaining why such RGA5_{HMA} homodimers do not occur in the RGA5_S/dSP-AVR1-CO39 complex (Fig. 1 and *SI Appendix, Figs. S4 and S5*).

Binding of AVR1-CO39 and AVR-Pia to RGA5_S Compete with RGA5_S Self-Interaction. Analytical gel filtration (GF) was performed to test the competition between RGA5_S/dSP-AVR1-CO39 complex formation and RGA5_S self-interaction. This showed that

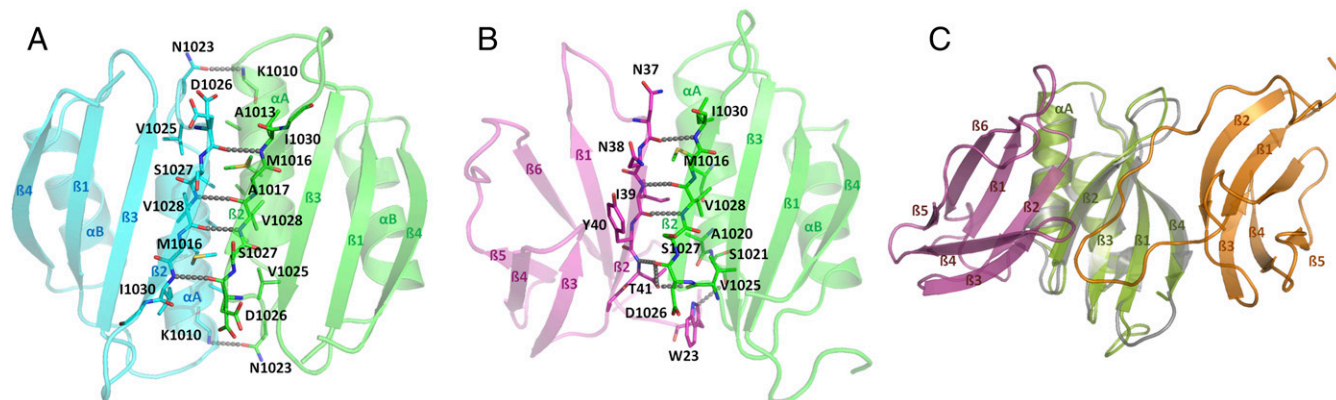


Fig. 1. AVR1-CO39 and AVR-PikD bind to opposite surfaces of the HMA-IDs. (*A*) The structures of the RGA5_{HMA} homodimer and (*B*) the RGA5_{HMA}/dSP-AVR1-CO39 complex are shown as cartoons with the RGA5_{HMA} molecules in cyan or green and the effector in magenta. Key residues in the binding interface are shown as sticks with carbon residues in the color of the cartoon. Hydrogen bonds are depicted as dashed lines. (*C*) Overlay of the structures of the RGA5_{HMA}/dSP-AVR1-CO39 (green and magenta, respectively) and the Pikp-1_{HMA}/dSP-AVR-PikD complexes (gray and orange, respectively) shown as cartoons.

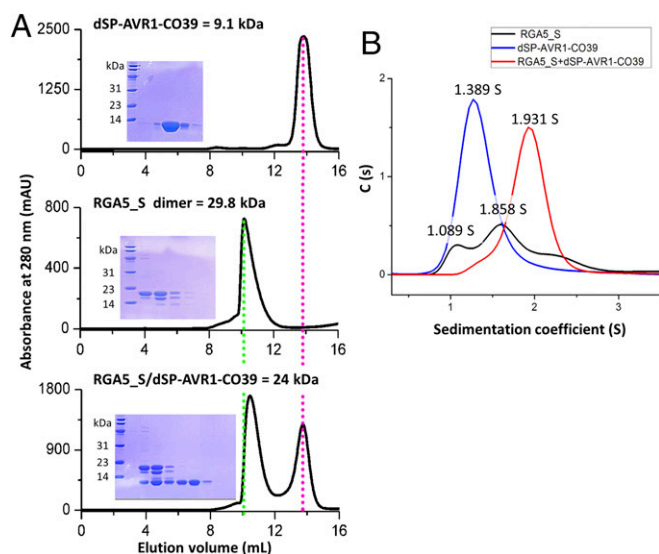


Fig. 2. AVR1-CO39 competes with RGA5_{HMA} self-interaction for RGA5_S-binding in vitro. (A) Gel filtration traces showing the retention volume and SDS/PAGE gels of relevant elution fractions of dSP-AVR1-CO39, RGA5_S, and a 1:1 mixture of both proteins. The molecular weight of the recombinant proteins is depicted on the *Top* of the panels. (B) Sedimentation coefficient distributions calculated from sedimentation velocity analytical ultracentrifugation with dSP-AVR1-CO39, RGA5_S, and a 1:1 mixture of both proteins. These GF (A) and SV-AUC (B) experiments were performed three times with similar results.

dSP-AVR1-CO39 is, as previously described, monomeric in solution (12), while RGA5_S forms homodimers (Fig. 2A). When dSP-AVR1-CO39 and RGA5_S were combined at a 1:1 ratio, two elution peaks with retention times indistinguishable from the dSP-AVR1-CO39 monomer peak and the RGA5_S dimer peak were detected (Fig. 2A). SDS/PAGE analysis of the elution fractions showed coelution of dSP-AVR1-CO39 with RGA5_S in the first peak, indicating formation of an RGA5_S/dSP-AVR1-CO39 heterocomplex with a 1:1 stoichiometry (Fig. 2A).

In sedimentation velocity analytical ultracentrifugation (SV-AUC), dSP-AVR1-CO39 occurred in one uniform state with a sedimentation coefficient of 1.389 S, while RGA5_S occurred in two main states: a small fraction (8.5%) with a low sedimentation coefficient (1.089 S) that contains likely monomeric RGA5_S and a major fraction (85%) with a sedimentation coefficient of 1.858 S that probably corresponds to the dimer (Fig. 2B). A small fraction of RGA5_S with even higher sedimentation coefficients corresponds potentially to higher-order oligomers. When dSP-AVR1-CO39 and RGA5_S were mixed, we detected only one predominant peak, whose sedimentation coefficient differs from those obtained with single proteins and that probably corresponds to the RGA5_S/dSP-AVR1-CO39 complex (Fig. 2B). This further confirms that RGA5_S forms homodimers in solution and that there is competition between RGA5_S/dSP-AVR1-CO39-binding and RGA5_S self-interaction.

GF and SV-AUC analysis were also performed with dSP-AVR-Pia and RGA5_S and showed that this MAX effector also competes with RGA5_S self-interaction, suggesting that the AVR-Pia-binding interface in RGA5_{HMA} overlaps with the α A- β 2 homodimerization surface (SI Appendix, Fig. S2 B and C).

ITC Measurements and NMR Titration Support the RGA5_S/dSP-AVR1-CO39 Structure and Validate the RGA5_{HMA}-Binding Surface in AVR1-CO39. The RGA5_S/dSP-AVR1-CO39 structure indicates that residues of RGA5_S outside the HMA domain are not involved in AVR1-CO39-binding since they are not resolved. To test this, the binding strength of dSP-AVR1-CO39 to a minimal RGA5_{HMA} construct (RGA5₉₉₁₋₁₀₇₂, SI Appendix, Fig. S1) was determined by

ITC. A K_d of 7.2 μ M, almost identical to the K_d for RGA5_S/dSP-AVR1-CO39 (5.4 μ M), was found (17), indicating that residues of RGA5_S outside of the HMA domain do not contribute to AVR1-CO39-binding (SI Appendix, Fig. S6).

To further investigate complex formation in solution, NMR titration experiments were performed where ^1H - ^{15}N -HSQC NMR spectra of ^{15}N -labeled dSP-AVR1-CO39 were recorded in the absence and presence of unlabeled RGA5_{HMA}. Mapping of the NH chemical shifts identified two major sequence stretches where most of the conformational rearrangements occur (Fig. 3 A–C): the first at the N terminus corresponds to the beginning of β 1 and the second to the β 1- β 2 turn and β 2. Together, these residues form the three main binding areas detected in the crystal structure of the RGA5_S/dSP-AVR1-CO39 complex (SI Appendix, Fig. S5). The low chemical shift differences in the rest of the molecule indicate that residues in the strands β 3, β 4, β 5, and β 6 are not perturbed and not involved in the interaction with RGA5_{HMA}. NMR titration therefore defines the same binding interface as the crystal structure.

Structure-Guided Mutations of Key Residues in the RGA5_{HMA}-Binding Surface of dSP-AVR1-CO39 Disrupt Complex Formation in Vitro and in Vivo. Based on the RGA5_{HMA}/dSP-AVR1-CO39 structure, we designed mutations in the HMA-binding surface of AVR1-CO39 to perturb complex formation: seven dSP-AVR1-CO39 mutants with substitutions in the main RGA5_{HMA}-binding area (W23S, K24A, K24E, W23A/K24A, T41G, T41A, and T41V), one mutation in the second area (N38A), two in the third area (D35A, N37G), and one double mutation targeting both the first and third area (K24A/D35A) (SI Appendix, Fig. S4). These mutants were tested in yeast two-hybrid assay for interaction with RGA5_L (Fig. 4A and SI Appendix, Fig. S1). A complete loss of interaction was observed for dSP-AVR1-CO39^{W23A/K24A} and dSP-AVR1-CO39^{T41G}, which both affect the main RGA5_{HMA}-binding area (Fig. 4A). A severe reduction of binding to RGA5_L was caused by the mutations K24A, K24E, T41A, T41V, and K24A/D35A, while the D35A and N37G mutations in the third binding area only mildly reduced the

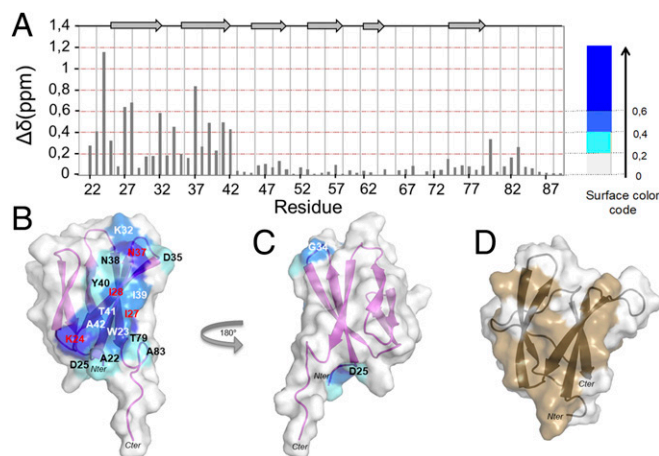


Fig. 3. AVR1-CO39 binds RGA5_{HMA} through the interface defined by the crystal structure of the complex. (A) Plot of the chemical shift differences ($\Delta\delta$ (ppm)) between unbound ($R = 0$) and RGA5_{HMA}-bound dSP-AVR1-CO39 ($R = 2$). Chemical shift differences were calculated as the Hamming distance (31), $\Delta\delta(\text{ppm}) = |\Delta\delta(^1\text{H})_{ij}| + 0.102 \times |\Delta\delta(^{15}\text{N})_{ij}|$, where $\Delta\delta(^1\text{H})_{ij}$ and $\Delta\delta(^{15}\text{N})_{ij}$ are the differences of the ^1H and ^{15}N chemical shifts at $R = 0$ and $R = 2$, respectively. (B) Structure of dSP-AVR1-CO39 showing the chemical shift differences from NMR titration with the following color code: surfaces of residues with $\Delta\delta(\text{ppm}) \geq 0.6$ in dark blue (residues in red letter), $0.6 > \Delta\delta(\text{ppm}) \geq 0.4$ in light blue (residues in white letters), and for $0.2 > \Delta\delta(\text{ppm}) \geq 0.4$ in cyan (residues in black letters). (C) 180° rotation of B. (D) Structure of the WT dSP-AVR1-CO39 with the RGA5_{HMA} interaction surface previously determined by NMR (14) in brown.

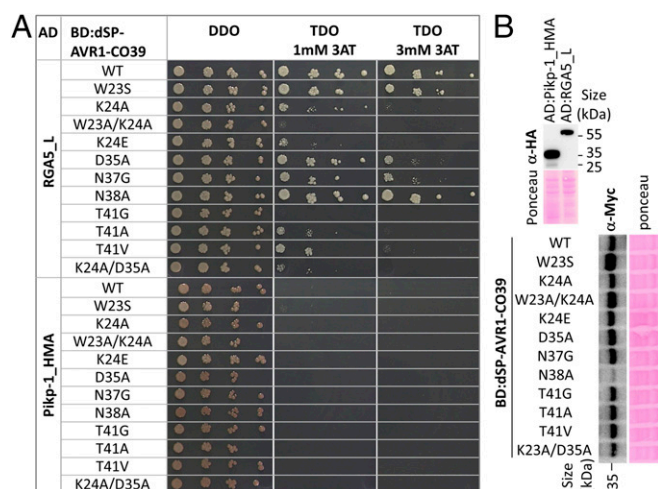


Fig. 4. Key residues within the HMA-binding surface of AVR1-CO39 are required for in vivo association with RGA5_L. (A) Interaction of RGA5_L (residues 883–1116, AD fusion) with dSP-AVR1-CO39 wild type and variants carrying point mutations in the HMA-binding surface (BD fusion) in yeast two-hybrid experiments. The HMA domain of Pk1p-1 (AD:Pk1p-1_{HMA}) was used as a control. Four dilutions of diploid yeast clones (1/1, 1/10, 1/100, 1/1000) were spotted on TDO medium (–Trp–Leu–His supplemented with 1 or 3 mM of 3-amino-1,2,4-triazole) to assay for interactions and on DDO (–Trp–Leu) to monitor proper growth. Pictures were taken after 5 d of growth. The experiment was performed three times with consistent results. (B) Yeast protein extracts were analyzed by immunoblotting using anti-myc or anti-HA antibodies.

interaction. The W23S and N38A mutations did not affect binding to RGA5_L. Pk1p-1_{HMA} was used as a control and showed no interactions. All Gal4 DNA-binding domain (BD)- or Gal4 activation domain (AD)-fused proteins were expressed according to immunoblotting (Fig. 4B).

To test the interaction of the dSP-AVR1-CO39 mutants with RGA5_S in vitro, recombinant proteins were produced (except K24E, T41A, and T41V). The dSP-AVR1-CO39^{W23A} and dSP-AVR1-CO39^{W23A/K24A/T41G} mutants were also expressed to further assess the contribution of binding area 1 to RGA5_{HMA}-binding. Soluble proteins were obtained for all mutants except K24A, D35A, N37G, and N38A and were analyzed by circular dichroism for proper folding (SI Appendix, Fig. S7A). ITC analyses showed that replacement of W23 by S or A only slightly reduced the affinity of dSP-AVR1-CO39 to RGA5_{HMA} (K_d s of 9.8 μ M and 11.4 μ M), while the double mutants W23A/K24A and K24A/D35A had drastically reduced affinities (Fig. 5).

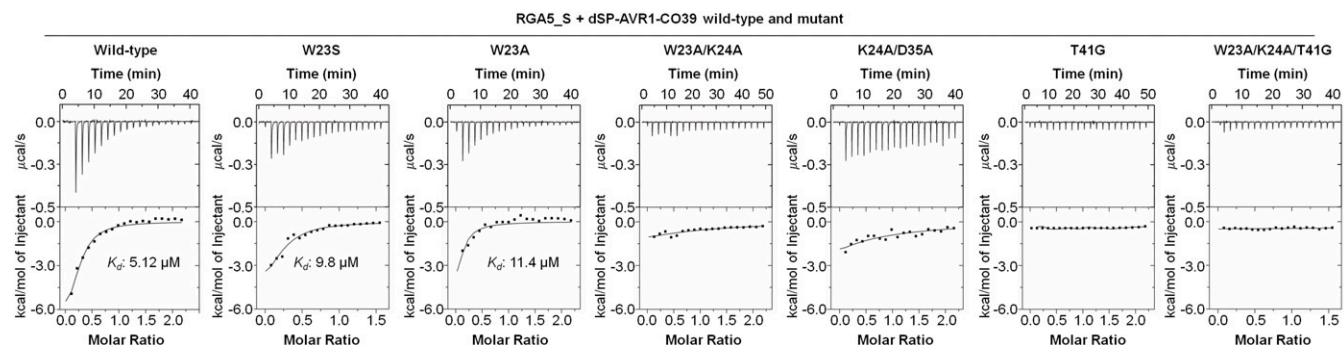


Fig. 5. In vitro binding assays demonstrate the importance of binding area 1 in the HMA-binding interface of AVR1-CO39 for RGA5_S/dSP-AVR1-CO39 complex formation. ITC curves for the titration of RGA5_S by dSP-AVR1-CO39 variants carrying point mutations in the HMA-binding surface. To ensure easy comparison, mutant proteins were used in the same concentration range as the AVR1-CO39 wild-type protein. Bottom shows the integrated heats of injection (squares) and the best fit (solid line) to a single-site-binding model. The assays were performed at least three times with similar results.

Replacement of T41 by G in dSP-AVR1-CO39^{T41G} or dSP-AVR1-CO39^{K24A/D35A/T41G} completely abolished binding.

Further investigation by GF confirmed that the dSP-AVR1-CO39 mutants carrying the W23A or W23S substitution still bind RGA5_S, while the T41G and W23A/K24A/T41G mutations abolished dSP-AVR1-CO39-binding to RGA5_S (SI Appendix, Fig. S8).

Since the T41G mutation so strongly affects RGA5_{HMA}-binding, dSP-AVR1-CO39^{T41G} and dSP-AVR1-CO39^{W23A/K24A/T41G} structures were reassigned using ¹⁵N-NOESY-HSQC and ¹⁵N-TOCSY-HSQC 3D NMR experiments. The chemical shift differences between the mutants and the wild-type dSP-AVR1-CO39 (SI Appendix, Fig. S7B and C) show that replacement of T41 by a glycine propagates to the residues close in space in particular I27 and I28 in β 1 that show strong chemical shift variation. Interresidue contacts observed in the ¹⁵N-NOESY-HSQC 3D NMR spectra indicate that while β -strands are essentially conserved, several interstrand long-range contacts involving residues in β 1 are missing in the mutants. For example, the contact between HN-I28 (β 1) and H α -T41 (β 2), found in the wild-type protein, is missing in both dSP-AVR1-CO39^{T41G} and dSP-AVR1-CO39^{W23A/K24A/T41G}. Since β 2 of dSP-AVR1-CO39 is central for RGA5_{HMA}-binding, the local distortion and probable flexibility at the C terminus of β 2 caused by the T41G mutation is believed to be the main factor that destabilizes the complex.

To determine whether dSP-AVR1-CO39 associates with RGA5 through the same interface in planta as in vitro and in yeast, we performed coimmunoprecipitation (Co-IP) experiments in *Nicotiana benthamiana* using YFP-tagged dSP-AVR1-CO39 mutants, and HA-tagged RGA5_L. HA:RGA5_L was coprecipitated with all YFP:dSP-AVR1-CO39 constructs except the T41G, W23A/K24A, K24E and W23S mutants (SI Appendix, Fig. S9). For T41G and W23A/K24A, this is consistent with the Y2H, ITC, and GF data. For the W23S mutant, loss of association with RGA5_L was not observed in other assays. The K24A/D35A mutation drastically reduced RGA5_{HMA} interaction in ITC and Y2H, but not in planta. As in the Y2H assay, the in planta association of RGA5_L with AVR1-CO39 was reduced in the T41A and D35A mutants and not affected by N37G or N38A.

These results show that key residues in the main RGA5_{HMA}-binding area of AVR1-CO39 (in particular, W23, K24, and T41) are crucial for the interaction with RGA5_{HMA}. Residues D35 and N37 within the second interaction area contribute weakly to RGA5_{HMA}-binding, while N38 in the third area does not play a major role.

RGA5_{HMA}/AVR1-CO39 Complex Formation Is Required for Resistance.

To evaluate whether binding of AVR1-CO39 to RGA5_{HMA} is required for immunity, transgenic *M. oryzae* isolates carrying AVR1-CO39 variants affected at various degrees in their interaction with

RGA5_{HMA} were generated together with positive (wild-type *AVR1-CO39*) and negative (*mRFP*) controls. Infection experiments with the susceptible rice variety Nipponbare (*pi-co39*) demonstrated that all transformants possess full virulence (*SI Appendix*, Fig. S10). Quantification of transgene expression by qRT-PCR showed that *AVR1-CO39* was expressed in the transgenic isolates (*SI Appendix*, Fig. S11). On the resistant Kitaake variety (*Pi-CO39*), only *AVR1-CO39*^{D35A} triggered immunity at the same level as wild-type *AVR1-CO39*, which is consistent with the low impact of this residue on interaction with RGA5_L in Y2H and Co-IP (Fig. 6). All other mutants induced disease symptoms ranging from small lesions, characteristic of mild recognition, to fully expanded disease lesions, identical to the virulent *mRFP* control isolate and indicating loss of effector recognition.

Overall, these data highlight an important role of the HMA-binding surface of *AVR1-CO39* in its recognition by the RGA5 immune receptor. However, *AVR1-CO39* recognition seems to be quite resilient to modifications in this interaction surface, as only the triple mutant, affected in three important residues of the main area, showed full virulence and complete loss of recognition.

Discussion

The identification of unconventional IDs in NLRs, as effector detector modules, was a major breakthrough and led to the development of the integrated decoy model for effector recognition (7, 21, 22). This model states that IDs are paralogous to and act as mimics of true effector target proteins and that the presence of pathogens is sensed either by effector-mediated modification of the ID or by its interaction with the effector. In the present study, we demonstrate that binding of the *M. oryzae* effector *AVR1-CO39* to the HMA ID of the rice NLR RGA5 contributes to effector recognition. Indeed, the binding strength of mutants of this MAX effector to the isolated RGA5_{HMA} ID is correlated with the level of immunity against *M. oryzae* elicited on a

resistant rice variety. Similar results have also been observed in the case of RGA5_{HMA}/*AVR-Pia* and P1kp-1_{HMA}/*AVR-PikD* (13–15). We therefore provide an additional demonstration of the important role of direct ID/effector interaction in the specific recognition of effectors.

By solving the RGA5_{HMA}/*AVR1-CO39* crystal structure and through NMR titration experiments, we found that the RGA5_{HMA}-binding surface in *AVR1-CO39* overlaps with the RGA5_{HMA}-binding surface in *AVR-Pia* (14). Indeed, as in *AVR1-CO39*, N-terminal residues of *AVR-Pia* located before β 1, as well as residues in β 2, mediate the interaction (14). However, in *AVR-Pia*, the HMA-binding surface is more extended than in *AVR1-CO39* and also involves the loop between β 2 and β 3 as well as the beginning of β 3 (Fig. 3D). GF and AUC data indicate that, like *AVR1-CO39*, *AVR-Pia* competes with RGA5_{HMA} self-interaction, suggesting that it also binds to the α A/ β 2 surface of the HMA. Further elucidation of this complex awaits a crystal structure of the RGA5_{HMA}/*AVR-Pia* complex that we could not yet obtain.

Our analysis reveals that the HMA IDs of P1kp-1 and RGA5, which both interact with structurally conserved MAX effectors and are important for NLR receptor activation, are structurally highly similar and self-interact through almost identical interfaces formed by α A and β 2. Besides, the formation of the RGA5_{HMA}/*AVR1-CO39* and P1kp-1_{HMA}/*AVR-PikD* complexes shows, at the molecular level, important similarities. In both complexes, binding involves antiparallel alignment of the β -sheet of the HMA with one of the two β -sheets of the MAX effector, resulting in one antiparallel continuous β -sheet in the hetero-complex. However, there are also fundamental differences. P1kp-1_{HMA}/*AVR-PikD* interaction relies very much on the N-terminal extension of *AVR-PikD* that is not part of the core MAX effector fold and makes strong and highly specific interactions with a defined area of the P1kp-1_{HMA} surface (13). Conversely, RGA5_{HMA}/*AVR1-CO39*-binding exclusively relies on interactions with the core MAX effector domain and does not involve the C-terminal unstructured extension of *AVR1-CO39*. In addition, in the two complexes, effector-binding is mediated by different surfaces of both the HMA-ID and the MAX effector. In the RGA5_{HMA}/*AVR1-CO39* complex, the β 2s of both partners are aligned, while in the P1kp-1_{HMA}/*AVR-PikD* complex, β 4 of the HMA is aligned with β 3 of the effector. Therefore, effector-binding involves different β -sheets of the effectors and occurs on opposite sides of the HMAs. *AVR1-CO39* disrupts the RGA5_{HMA} homodimer by competing for β 2/ α A-binding, while in the P1kp-1_{HMA}/*AVR-PikD* complex, the β 2/ α A surface is free for self-interaction and, consequently, P1kp-1_{HMA} occurs as a dimer in the P1kp-1_{HMA}/*AVR-PikD* crystal structure. The fact that HMA IDs in different NLRs interact through completely different interfaces with sequence-unrelated but structurally conserved MAX effectors has major implications. It suggests that these interactions do not rely on conservation but evolved independently by convergence of the different MAX effectors to the same type of plant target proteins.

An exciting perspective of this finding, from an applied point of view, is that it may be possible to engineer these HMA IDs for extended recognition specificity, allowing them to recognize both *AVR1-CO39* and *AVR-PikD*. Indeed, introduction of the additional binding site by changing the relevant surface-exposed residues should, in each case, not interfere with the original binding site located at the opposing site of the HMA.

The relevance of HMA self-interactions for RGA5 function is unclear, and it is unknown whether it occurs in the context of the full length RGA5 receptor. RGA5 forms heterocomplexes of unknown stoichiometry with RGA4, a constitutively active immune activator that is repressed by RGA5 in the resting state and triggers immunity upon RGA5-mediated *AVR1-CO39* recognition (10, 16). RGA5 was also shown to form homocomplexes in planta, suggesting that the RGA4/RGA5 inactive complex may incorporate more than one RGA5 protein (16). In this context, RGA5_{HMA} self-interaction could be relevant for RGA4 repression, and a switch to “activated state” might be triggered

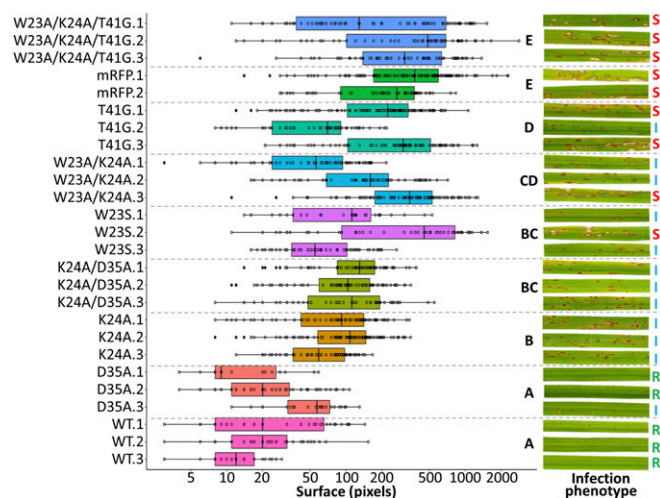


Fig. 6. Key residues within the HMA-binding surface of *AVR1-CO39* are important for RGA5-mediated effector recognition in rice. Transgenic *M. oryzae* isolates carrying wild-type *AVR1-CO39* (WT), *AVR1-CO39* variants affected in the HMA-binding surface or *mRFP* (control), were spray-inoculated on plants of the rice cultivar Kitaake (*Pi-CO39*). Three independent transgenic isolates were generated for each construct (two isolates for *mRFP*). Leaves from 10 different plants for each isolate were scanned 7 d after inoculation, and areas of disease lesions were measured and plotted. The boxes represent the first quartile, median, and third quartile. Difference of lesion areas was assessed by an ANOVA followed by a Tukey HSD test. Groups with the same letter (A to E) are not significantly different at level 0.05. Representative disease phenotypes are shown (Right) for each strain (I, intermediate; R, resistant; S, susceptible). The experiment was performed two times with similar results.

upon effector-mediated disruption of the RGA5_{HMA} dimer. Alternatively, RGA5_{HMA} self-interaction may not be relevant for RGA5 activity, either because the α A/ β 2 HMA surface is unoccupied in the inactive receptor complex or because it is engaged in other intra- or intermolecular interactions, potentially required for RGA4 repression. In this case, AVR1-CO39-binding to RGA5_{HMA} could disrupt such interactions, resulting in conformational changes that would lead to RGA4 de-repression and immune activation. To test these hypotheses, it will be important to identify RGA5_{HMA} mutations that uncouple self-interaction from AVR1-CO39-binding and to further search for alternative inter- or intramolecular interactions of the HMA ID.

Virulence targets of AVR1-CO39, AVR-Pia, or AVR-PikD have not been described yet. However, the integrated decoy model suggests that small HMA proteins (sHMAs), paralogous to Pk1p-1_{HMA} and RGA5_{HMA}, are among them. The differential interaction of the two MAX effectors with their cognate HMA IDs suggests that AVR1-CO39 and AVR-PikD evolved independently toward these targets and target them in different manners. Whether AVR1-CO39 acts by competing with sHMA homo- or heterodimerization through the α A- β 2 surface remains to be investigated since homocomplex formation of plant sHMAs has not been reported, and knowledge on sHMAs, which represent in plants a huge and rapidly evolving protein family, is extremely limited (23, 24). The importance of sHMAs for plant-pathogen interactions is strongly supported by the finding that the rice sHMA Pi21 is required for full susceptibility to the blast fungus (25). Therefore, there is a strong need to test whether sHMA proteins are virulence targets of MAX effectors and to elucidate their molecular and cellular functions.

Our finding that the interaction of AVR1-CO39 and AVR-PikD with HMA domains is presumably a consequence of convergent evolution toward similar host proteins suggests that MAX effectors potentially target other host proteins besides sHMAs. This hypothesis is supported by work on the *M. oryzae* MAX effector AvrPiz-t, for which five different host target proteins have been described (i.e., two E3 ubiquitin ligases, a K⁺ transporter, a b-ZIP transcription factor, and a nucleoporin protein Nup98) (26–30). These observations raise a fundamental

question: How can these distinct MAX effectors target such a wide diversity of host proteins? Our study suggests that the molecular mechanism by which AVR1-CO39 and AVR-PikD interact with HMA domains is conserved. In both cases, binding relies on antiparallel alignment of one β -sheet of the MAX effector with the β -sheet of the HMA domain. This involves either β 2 or β 3 of the MAX effector, engaged in multiple backbone and side-chain interactions with the target β -structure. In addition, variable additional interactions by residues in β 1 or from sequences outside the core MAX effector fold, such as the short and long N-terminal extensions of AVR1-CO39 and AVR-PikD, respectively, stabilize the interaction. Therefore, we hypothesize that, in the structurally conserved MAX effector family, the structural determinants and molecular mechanism of host target protein-binding are conserved rather than the host target proteins themselves. This establishes a framework for the understanding of the evolution and diversification of fungal effectors that can be tested by identifying additional host target proteins of MAX effectors and elucidating the structural details of MAX effector/target protein-binding.

Methods

Experimental procedures can be found in *SI Appendix*, including molecular biology, protein expression in *Escherichia coli* and *N. benthamiana*, yeast two-hybrid analysis, biochemical and physical characterization of proteins, and their complexes and infection assays.

ACKNOWLEDGMENTS. The work was supported by the National Natural Science Foundation of China (NSFC) (Grant 31571990); the Agence Nationale de la Recherche (ANR) (Grant ANR-15-CE20-0007); the National Key Research and Development Program (Grant 2016YFD0300700) from the Ministry of Science and Technology, China; the 111 Project (B13006) and the Program for Changjiang Scholars and Innovative Research Team in University (PCSIRT) Project (IRT1042) from the Ministry of Education, China; the Project for Extramural Scientists of State Key Laboratory of Agrobiotechnology (Grant 2017SKLAB7-9); and the French Infrastructure for Integrated Structural Biology (Grant ANR-10-INBS-0005). This work also benefited from interactions promoted by COST (European Cooperation in Science and Technology Program) Action FA 1208.

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